

‘Future-proofing’ blood processing for measurement of circulating microRNAs in samples from biobanks and prospective clinical trials

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Abstract

Background: Quantifying circulating nucleic-acids is an important new approach to cancer diagnosis/monitoring.

Methods: We compared the suitability of serum versus plasma for measuring microRNAs using RT-qPCR and assessed how pre-analytical variables that can affect circulating-tumor-DNA (ctDNA) quantification in plasma also influence microRNA levels.

Results: Across 62 blood-derived specimens, plasma samples in EDTA, Streck-DNA-plasma and Streck-RNA-plasma tubes showed significantly higher Ct values for multiple housekeeping microRNAs, compared with serum samples. For the EDTA-plasma tubes, this difference was only seen when including the high-speed centrifugation protocol used to optimise ctDNA extraction. In plasma samples derived from blood stored at room temperature for up to 14 days (conditions that typically apply to samples processed for biobanking), levels of endogenous housekeeping microRNAs gradually increased, in parallel with the hemolysis-marker hsa-miR-451a, consistent with release from blood cells/platelets. It was necessary to normalize levels of the housekeeping microRNAs to those of hsa-miR-451a, in order to obtain the stable values needed for referencing test microRNA levels.

Conclusions: Our data indicate that plasma samples prepared for ctDNA extraction are suboptimal for microRNA quantification and require the incorporation of multiple data normalization steps. For prospective studies designed to measure both microRNAs and ctDNA, the most suitable approach would be to obtain both serum (for microRNAs) and plasma (for ctDNA). If only plasma can be collected, we recommend an initial low-speed centrifugation step, followed by aliquoting the supernatant into parallel samples, one for direct microRNA quantification, and the other for a further high-speed centrifugation step to optimize ctDNA retrieval.

Impact: These recommendations will help ‘future-proof’ clinical studies in which quantification of circulating microRNAs is a component.

Introduction

New approaches to cancer diagnosis and/or monitoring are based on detecting nucleic acids in blood, including circulating tumor DNA (ctDNA) (1-6) and microRNAs (7,8). Studies using ctDNA have almost exclusively been based on plasma, due to the low yield of ctDNA retrieved from serum (9,10). Stability of ctDNA in plasma is improved using preservatives, for example those in Streck tubes (11-13). However, it is clear that multiple pre-analytical variables must be considered in prospective ctDNA-based clinical trials, including time to plasma isolation and the types of preservative used (14).

Detection of microRNAs is particularly attractive for many non-epithelial and/or pediatric cancers that have a low mutational prevalence (15,16). In testicular malignant germ cell tumors (GCTs), for example, mutations occur in less than half of all cases (17). We previously identified that microRNAs from the miR-371-373 and miR-302/367 clusters were co-ordinately elevated in the serum at the time of malignant GCT diagnosis (18,19) and were also sensitive indicators of disease relapse (20). Subsequent studies, from multiple groups, have all focussed on serum samples and have confirmed the clinical importance of this novel approach to malignant GCT diagnosis (20-30). In consequence, collection of biospecimens including blood for further evaluation of microRNA detection has been embedded in multiple prospective clinical trials across the spectrum of GCTs.

As future studies of circulating nucleic acids are likely to compare or combine microRNAs and ctDNAs for tumor diagnosis/monitoring (31), we sought to address whether plasma would also be a suitable medium for detecting circulating tumor-derived microRNAs. If so, this would allow combinatorial analysis of microRNAs and ctDNAs from the same sample.

We also assessed whether pre-analytical factors that can affect ctDNA analysis in plasma samples would also affect microRNA levels in such samples. We compared serum obtained using gel separator tubes with plasma derived from EDTA, Streck DNA and Streck RNA tubes. For the EDTA tubes, we used two-step centrifugation protocols optimized for ctDNA extraction from plasma (32-34). We measured circulating levels of established housekeeping microRNAs and assessed effects on microRNA levels of tube type, time to processing, centrifugation speed and hemolysis. Based on our findings, we recommend protocols for microRNA detection that are optimised for blood samples from biobanks and biological studies linked to clinical trials.

Materials and Methods

Patient samples

In total, we analyzed 62 blood-derived specimens. First, we undertook a comparison study, in which we comprehensively analyzed 27 specimens collected from a cohort of adult male patients (n=7) recruited from a single hospital, via the Cambridge Urology Translational Research and Clinical Trials Biorepository (CUTRACT; cases CUB_001 – CUB_007). This cohort comprised a group with testicular malignant GCTs (n=5) and a control group with non-GCT testicular malignancy (n=2; one relapsed acute myeloid leukemia and one B-cell lymphoma). Second, we performed a time-course study of 26 samples from healthy adult male subjects (n=2), plus nine specimens from a patient with testicular Leydig cell tumor (CUB_008). Details of all specimens, including clinico-pathological information, are provided in Supplementary Table S1. All samples were collected with Local Research Ethics Committee approval (CUB samples reference 03/018; other samples reference 01/128) and informed consent.

Sample collection and processing

For the comparison study, blood samples were collected immediately prior to orchiectomy. The tubes used were: 4.9ml Sarstedt S-Monovette Z-Gel (gel separator serum; ‘GSS’), 9ml Sarstedt S-Monovette K3E (EDTA plasma; ‘EP’), 10ml Streck Cell-Free DNA BCT (Streck DNA plasma; ‘SDP’) and 10ml Streck Cell-Free RNA BCT (Streck RNA plasma; ‘SRP’). Six patients provided samples in all four tubes, while one patient (CUB_003) provided samples in three tubes, as an SDP tube was unavailable at the time of patient recruitment. All specimens were centrifuged and processed at room temperature within two hours of

venepuncture, according to the recommendations of the manufacturers of each tube. There was no difference in processing times (average or range) for the different tubes in the comparison study. Details of the processing are summarized in Supplementary Table S2. In brief, for GSS tubes, the blood was allowed to clot for 30 minutes (min), then centrifuged at 3,000g for 10min and the separated serum removed from above the gel layer. For EP tubes, the blood was centrifuged at 1,600g for 10min, then the separated plasma was aliquoted and subjected to a secondary spin of 14,400g for 10min, as per standard protocol conditions for preparing plasma for optimal ctDNA analysis (32-34). For SDP and SRP tubes, the blood was initially centrifuged at 300g for 20min and the separated plasma aliquoted into fresh tubes, prior to a secondary spin of 5,000g for 10min. For all samples, the final supernatants were aliquoted into fresh tubes, then stored immediately at -80°C until further analysis.

In the time-course study, we used 2ml SDP tubes, in which levels of RNA recovery and endogenous microRNAs were highly comparable to those in 10ml SDP tubes. Blood samples were collected into 1.2ml EP and 2ml SDP tubes, then stored at room temperature for 0, 2, 4, 7, 10 and 14 days before being processed as above. The samples from one subject (patient CUB_008) were also used to study the effects of centrifugation speed on microRNA recovery. Of two 1.2ml EP tube samples from this patient, one (EP #1) underwent only the first low-speed centrifugation (1,600g) only, whereas the other (EP #2) also underwent the second high-speed centrifugation (14,400g).

RNA extraction

RNA was isolated from 200µl of thawed serum or plasma using the miRNeasy serum/plasma kit (Qiagen, Crawley, UK), incorporating MS2 carrier RNA (Roche, Welwyn Garden City,

UK) and the exogenous non-human microRNA spike-in, cel-miR-39-3p, as described (20). RNA was eluted in 100µl of nuclease-free water. For the comparison study, 750µl of QIAzol per sample was used. To assess whether the presence of proteins/nucleases within the biospecimens affected RNA retrieval, a subset of samples underwent a further RNA extraction that included a proteinase K digestion step. For this, 15µl proteinase K (minimum concentration >600mAU/ml; Qiagen) was added to 200µl of each sample, then incubated at 60°C for 60 minutes. Subsequently, 750µl of the QIAzol/MS2/cel-miR-39-3p mix was added, following which the protocol proceeded as described (20). The same protocols were used in the time-course study, except that the volume of QIAzol per sample was increased to 1000µl, following an update to the manufacturer's recommendations.

We also assessed two alternative approaches to RNA extraction from SRP tubes, namely the Plasma/Serum RNA Purification Mini Kit (Norgen BioTek Corp, Thorold, ON, Canada) and the mirVana PARIS isolation kit (Ambion, Warrington, UK). For the Norgen kit, 200µl of circulating sample was used, with the addition of cel-miR-39-3p/MS2 as above, to the lysis buffer A. RNA was eluted using 50µl of nuclease-free water. The standard protocol for the Norgen plasma/serum kit was also adjusted to include a proteinase K digestion step, where 20µl of the proteinase K solution described above was added to lysis buffer A, mixed with the sample and incubated for 60min at 60°C. For the mirVana kit, 400µl of circulating sample was used, with the addition of 5.6×10^8 copies of cel-miR-39-3p spike-in and MS2 carrier RNA to the denaturing solution. In total, 80µl eluate was obtained.

RT-qPCR analysis of microRNA levels

In all experiments, levels of circulating microRNAs were detected using our singleplex RT-qPCR method, as described (20). In short, 5µl of eluted RNA was reverse transcribed using the TaqMan miRNA reverse transcription kit (Life Technologies, Paisley, UK), using the microRNA-specific stem-loop primer from the relevant TaqMan microRNA assay kit (Life Technologies). The final volume of 15µl for each reaction then underwent reverse transcription using a GeneAmp PCR System 9700 (Applied Biosystems, Warrington, UK) at 16°C for 30 min, 42°C for 30 min, followed by a final step of 85°C for 5min. A singleplex final PCR was then performed, as per the manufacturer's instructions, on a Mastercycler ep_gradient/S realplex (Eppendorf, Stevenage, UK) at 95°C for 10min, followed by 45 cycles of 95°C for 15s and 60°C for one minute, as described (20).

The Ct threshold on the PCR machine was set manually to 2,000 fluorescence units across all PCR plates (20). To exclude non-specific amplification, a no-template control (NTC) was also run for each assay. None of the test samples had expression levels within 2 Ct values of the relevant NTC samples. Samples with Ct values ≥ 40 , or those where the Ct threshold was not reached, were considered non-expressing and arbitrarily assigned a Ct value of 40. The standard deviations (SD) of Ct values for each set of technical triplicate RT-qPCR reactions were calculated. We compared Ct values and SD values between the different sample types. We defined Ct values for sample sets as being *suboptimal* if mean microRNA expression values were ≥ 2 Cts greater than the set with the lowest Ct value (i.e. expression levels were ≥ 4 -fold lower). We also defined SD values for sample sets as being *suboptimal* if they were ≥ 2 . For both Ct and SD assessment, samples that were not suboptimal were deemed to be *satisfactory*.

We quantified levels of six microRNAs, namely the exogenous spike-in microRNA cel-miR-39-3p (20) (added to each sample immediately prior to RNA extraction); the hemolysis-dependent hsa-miR-451a (35); and the endogenous human reference microRNAs hsa-miR-23a-3p, hsa-miR-30b-5p, hsa-miR-30c-5p and hsa-miR-191-5p, which were previously shown to be readily detectable in both serum and plasma (36,37). In addition, for the time-course study, we quantified levels of three further endogenous microRNAs, again using singleplex RT-qPCR, namely: hsa-miR-130b-3p and hsa-miR-146a-5p, previously shown to be present at stable levels in samples regardless of the degree of hemolysis (38); and hsa-miR-26a-5p, which had also been shown to be stable within the circulation (37).

Hemolysis assessment

In addition to visual inspection of each serum or plasma sample, hemolysis was assessed using two methods (8). First, spectrophotometric analysis of the absorbance of free hemoglobin at 414 nm (A_{414}) (38,39) was performed using a NanoDrop ND-1000 Spectrophotometer (NanoDrop Products, Wilmington, DE, US). Absorbance was tested in triplicate and the mean value recorded. Samples were classified as being hemolyzed if A_{414} values were >0.2 arbitrary units, as described (38,39). Second, using the RNA extracted from the sample, hsa-miR-451a Ct (35) and delta Ct (hsa-miR-23a-3p minus hsa-miR-451a) hemolysis values were calculated using singleplex RT-qPCR, as described (20). Delta Ct values ≥ 8 were considered to indicate hemolysis (20).

Statistical analysis

Statistics were performed using GraphPad Prism 6 software (San Diego, California, US). Unless otherwise stated, analyses were performed using either a two-tailed paired t-test (for

matched samples) or a two-tailed unpaired t-test with Welch's correction for unequal variance (for unmatched samples). *P*-values <0.05 were considered significant.

Results

Comparison study of microRNA levels in serum and plasma samples from different blood collection tubes

In the initial comparison study, we assessed levels of six microRNAs in serum samples prepared in GSS tubes, versus plasma samples prepared in EP, SDP or SRP tubes. Levels of each microRNA were quantified in each sample using technical triplicate RT-qPCR reactions. The SD values for each set of triplicate results were very low for all microRNAs across the GSS, EP and SDP samples. In contrast, the SD values were much higher for the samples in SRP tubes, particularly for hsa-miR-30b-5p and hsa-miR-23a-3p (Supplementary Figure S1).

Levels of each individual microRNA in samples from each tube type are shown in Figure 1.

In summary:

- For the exogenous normalization microRNA cel-miR-39-3p, Ct values and SDs were satisfactory and similar in the GSS, EP and SDP groups (Figure 1, Table 2), indicating acceptable RNA extraction and RT-qPCR efficiency. In contrast, these parameters were suboptimal in the samples in SRP tubes (Table 1), where Ct values were significantly greater than in the EP group (Supplementary Table S3).
- For the endogenous normalization microRNAs hsa-miR-30b-5p, hsa-miR-30c-5p, hsa-miR-191-5p and hsa-miR-23a-3p, Ct values were satisfactory only for serum samples in GSS tubes (Figure 1, Table 1). For the EP and SDP plasma samples, Ct values were suboptimal and significantly higher than in the samples in GSS tubes (Supplementary Table S3). For all four of the normalization microRNAs, the SRP

samples showed the highest Ct values, which were significantly greater than in the GSS and EP groups (Supplementary Table S3).

- For the hemolysis-associated microRNA hsa-miR-451a, Ct values were satisfactory for the GSS, EP and SDP groups (Figure 1, Table 1), but were again suboptimal in the SRP samples, where they were significantly higher than in each of the other sample groups (Supplementary Table S3).

We next compared mean Ct values for different microRNAs in each of the four tube types (Supplementary Figure S2). Variations in levels of cel-miR-39-3p and those of hsa-miR-30b-5p (Supplementary Figure S2) were lowest for samples in GSS tubes, which showed very tight clustering of Ct values. There was moderate clustering for the EP and SDP samples, but very poor clustering for the SRP samples (Supplementary Figure S2). Identical findings were also seen for hsa-miR-30c-5p, hsa-miR-191-5p and hsa-miR-23a-3p with cel-miR-39-3p. These data indicated that the EP, SDP and SRP tubes resulted in additional technical variation, when compared with GSS tubes.

We attempted to develop a method that would allow satisfactory RNA extraction and RT-qPCR analysis of microRNA levels in plasma from SRP tubes. . Extracting RNA from SRP tubes using the Norgen plasma/serum RNA purification kit reduced variability in levels of all six microRNAs, compared with RNA extracted from SRP tubes using the Qiagen miRNeasy serum/plasma kit (Supplementary Figure S3). However, mean Ct values of all six microRNAs in such SRP/Norgen samples remained ≥ 2 higher than those obtained with RNA extracted from GSS tubes using the Qiagen miRNeasy kit. Indeed, for the exogenous normalization microRNA cel-miR-39-3p, levels in the SRP/Norgen samples were almost 5 Ct

values greater than in the GSS/Qiagen samples. As the eluate from the former was 50 μ l and the latter 100 μ l, with 5 μ l from each being used for PCR, this amounted to ~58-fold less efficient RNA recovery from the original 200 μ l serum or plasma samples. Such differences in microRNA recovery using the Norgen protocol were not improved by modifications such as the incorporation of a proteinase K digestion step . When RNA was extracted from SRP tubes using the mirVana PARIS kit, expression levels of each of the six microRNAs tested (i.e. those listed in Table 1) were also suboptimal. This large reduction in the efficiency of RNA recovery would be expected to limit the overall sensitivity of a PCR assay for detecting test microRNA biomarkers (particularly those in low abundance), when compared with a serum-based approach. As we were unable to resolve the limitations of SRP tubes for quantifying circulating microRNAs, we did not include them in our subsequent analyses.

Time-course study of the effects of blood storage on microRNA levels in plasma

We studied the effects on microRNA quantification of sample storage in SDP tubes at room temperature for up to 14 days, which is the recommended time limit for sample preservation prior to plasma extraction for ctDNA quantification. We compared our findings with plasma samples stored at room temperature for the same time periods in EP tubes. In total, we analyzed 26 separate samples from two healthy subjects, comparing 14 samples in SDP tubes with 12 in EP tubes (Supplementary Table S1).

Levels of hemolysis were lower in the SDP samples compared with the EP samples at matched time-points, as indicated by sample inspection (Figure 2A) and by spectrophotometric measurement of free hemoglobin absorption at 414nm (A_{414}) (Figure 2B). Using the standard A_{414} threshold of >0.2 arbitrary units (38,39), the EP samples were

hemolyzed within 1 hour of collection (i.e. the d0 samples). They showed an abrupt increase in hemolysis from d4, consistently reaching the upper limit of detection threshold by d14. In contrast, hemolysis was not consistently present in the SDP samples until d7. Assessment of hemolysis using microRNA delta Ct values (hsa-miR-23a-3p – hsa-miR-451a) indicated hemolysis at almost all time-points in both sample types, albeit with occasional variation in the values observed (Figure 2C).

Next, we investigated changes in levels of individual microRNAs in plasma obtained from the SDP versus the EP tubes over the 14 day time-course (20). As cel-miR-39-3p was spiked-in immediately prior to RNA extraction for each time-point, it was not surprising that we saw no difference in levels of this microRNA (Figure 3A). In contrast, levels of all six endogenous housekeeping microRNAs tested [hsa-miR-30b-5p, hsa-miR-30c-5p, hsa-miR-191-5p (Figure 3); hsa-miR-26a-5p, hsa-miR-130b-3p, hsa-miR-146a-5p (Supplementary Figure S4)] all showed similar gradual reductions in Ct values over the time-course, indicating gradual increases in expression levels in the blood from which the plasma samples were derived. Similar changes were seen in the red blood cell derived microRNA hsa-miR-451a, suggesting that the elevations in endogenous housekeeping microRNAs were related to hemolysis. Indeed, normalization of levels of the six endogenous housekeeping microRNAs to those of hsa-miR-451a resulted in stabilization of microRNA measurements over the time-course (Figure 3 and Supplementary Figure S4). These findings were confirmed in a further set of SDP samples obtained from a patient with testicular Leydig cell tumor (CUB_008; Supplementary Table S1, Supplementary Figure S5).

Comparison of methods of hemolysis assessment in plasma samples

When quantifying test microRNAs, it is important also to measure hemolysis, in order to exclude any confounding effects of microRNAs released from blood cells (40). We investigated whether previously established parameters for measuring hemolysis in serum using microRNA quantification (8,20,41) could also be applied to the plasma samples in our study. The established delta Ct (hsa-miR-23a-3p - hsa-miR-451a) hemolysis method was applied to the samples from the comparison study in which there was no evidence of hemolysis by spectrophotometry (n=20) (Figure 4A). A delta Ct value indicating hemolysis (delta Ct >8) was seen in only one of seven GSS serum samples but in six of seven EP and five of six SDP plasma samples (Figure 4B). The false positive results in the plasma samples were attributable to significantly lower detection levels (higher Ct values) of the endogenous housekeeping gene hsa-miR-23a-3p in the EP and SDP samples, compared with the GSS samples ($p=0.018$ and $p=0.0002$, respectively) (Figure 1E, Supplementary Table S3).

When analysing 32 available EP and SDP samples from across the comparison and time-course studies, only 20 (62%) were classified as hemolyzed by spectrophotometry (Figures 4C/D), whereas 30 (94%) were classified as hemolyzed by using the delta Ct value (Figure 4D). The samples that were classified as hemolyzed by spectrophotometry showed higher delta Ct values ($p=0.009$, unpaired t-test; Figure 4D) and lower raw Ct values for hsa-miR-451a ($p<0.0001$, unpaired t-test; Figure 4C), compared with the samples that were classified as non-hemolyzed by spectrophotometry. The A_{414} absorbance values strongly fitted a non-linear association with the raw hsa-miR-451a Ct values ($R^2=0.818$; Figure 4E). However, they only weakly fitted a non-linear association with the delta Ct (hsa-miR-23a-3p – hsa-miR-451a) values ($R^2=0.224$; Figure 4F), in keeping with the limited accuracy of hsa-miR-23a-3p quantification in the plasma samples. The dynamic range of the A_{414} absorbance values was greater than those of the raw hsa-miR-451a values (Figure 4E) and delta Ct (hsa-

miR-23a-3p – hsa-miR-451a) values (Figure 4F), further indicating the limited utility of microRNA quantification for assessing hemolysis in the plasma samples.

Effect of centrifugation speed on microRNA levels in plasma

We sought to determine whether the differences observed between the serum and plasma samples were due to inherent features of the sample types or to the different processing protocols used. We examined the effect of centrifugation speed on microRNA recovery from plasma, using samples extracted from EP tubes, in comparison with serum samples prepared from GSS tubes. We first used samples from patient CUB_008, to compare the effect of a single low-speed centrifugation step (1,600g; EP #1) with the standard dual centrifugation that is typically used when preparing plasma for ctDNA analysis (1,600g then 14,400g; EP #2). The mean expression level of five microRNAs (namely hsa-miR-30b-5p, -30c-5p, -191-5p, -23a-3p and -451a) was significantly lower (higher Ct values) in the EP #2 sample, compared with the EP #1 ($p=0.03$) and the GSS sample ($p=0.01$) (Figure 5A). When analyzing each microRNA individually, levels were also lower in the EP #2 sample compared with the EP #1 and GSS samples (Figure 5B). There were no differences between the EP #1 and GSS samples in the mean expression levels (Figure 5A), nor in the levels of each microRNA individually (Figure 5B).

We extended these findings by interrogating data from a published global microRNA profiling study of EP samples (from breast cancer patients) that had been centrifuged at various speeds (42). For the 106 microRNAs with Ct values <35 [listed in (42)], mean levels were significantly lower in EP samples that underwent high-speed centrifugation (10,000g), compared with EP samples that underwent low-speed centrifugations (either 2,000g or

1,000g; $p < 0.001$ for both comparisons) (Figure 5C). There was no significant difference in mean microRNA expression levels between the 1,000g and 2,000g centrifugation samples (Figure 5C). Levels of individual microRNAs were again lower (higher Ct values) in the samples that had received high-speed centrifugation, compared with the low-speed samples (Figure 5D). The differences in expression levels were particularly large for the three endogenous housekeeping genes hsa-miR-30b-5p, hsa-miR-30c-5p, hsa-miR-191-5p, but were also seen for hsa-miR-451a (data on hsa-miR-23a-3p levels were not available). Across the 106 microRNAs, high-speed centrifugation had a highly unpredictable effect on microRNA levels compared with low-speed centrifugation (Figure 5E). Ct values were almost always higher in the high-speed samples, albeit with rare exceptions (Figure 5E). The difference in expression levels ranged from zero to >8 Ct values, with a modal difference of 2-4 Ct values (Figure 5F).

Discussion

Biospecimens collected by tissue banks increasingly include blood-derived samples for quantification of circulating nucleic acids. Most effort to date has focussed on studying pre-analytical variables that affect ctDNA quantification (11-14). Here, we sought to establish how such variables affect microRNA levels and to generate a recommended protocol for collecting and processing blood-derived biospecimens, in order to maximise the yield of recovered microRNAs. Our comparison study demonstrated that serum samples were better suited for microRNA studies than the plasma samples tested, as the latter showed greater technical variation, with consistently lower levels of endogenous housekeeping microRNAs. The differences we observed were not attributable to the method used for microRNA extraction, as the Qiagen miRNeasy serum/plasma kit was independently demonstrated to provide the highest yield of microRNAs from plasma samples (42). The difference between serum and plasma from EP tubes (Figure 1) is likely to relate to the two-spin processing used for preparing the latter, a method that is typically adopted to optimize ctDNA extraction. Samples processed from EP tubes with a low-speed centrifugation step only did not show significant differences in microRNA levels compared with serum samples (Figure 5A/B), indicating that plasma is not necessarily an inferior substrate for microRNA extraction, if appropriate protocols are adopted. We found Streck RNA (SRP) tubes to be unsuitable for microRNA studies, despite using various RNA extraction methods and protocol modifications, including the incorporation of a proteinase K digestion step.

The uniform clotting process inherent in preparing serum removes a large proportion of protein from a whole blood sample. One practical benefit of this is that the layer of protein obtained when extracting RNA from serum is relatively small. While endogenous

microRNAs are released into the serum during the clotting process, for example from platelets, the overall pipeline involved in serum processing is straightforward, and most importantly, provides very similar levels of endogenous housekeeping microRNAs across samples. As plasma lacks the clotting step, the protein layer obtained during RNA extraction is large, leading to smaller yields of supernatant. This may necessitate repeated sample processing in order to obtain adequate volumes of eluate for microRNA quantification, potentially exacerbating the technical variations in measuring microRNA levels in plasma.

A further consideration when evaluating blood-derived biospecimens is the stability of microRNAs during prolonged storage of unprocessed blood. Such conditions typically apply to the blood specimens from which plasma samples are obtained for biobanking, for example those from a multi-center clinical trial. In plasma samples derived from blood stored over a period of up to 14 days at room temperature, we observed increases in levels of the endogenous housekeeping microRNAs hsa-miR-30b-5p, hsa-miR-30c-5p and hsa-miR-191-5p. These were associated with parallel changes in the hemolysis marker hsa-miR-451a, suggesting that the microRNAs were likely to have been released into the plasma from red blood cells and/or platelets, either through active shedding or passively as a result of hemolysis. As a result, stabilization of the levels of housekeeping microRNAs in plasma samples over the 14 day time-course required correction for levels of hsa-miR-451a.

An additional consequence of the variations in levels of housekeeping microRNAs in the plasma samples studied is that hemolysis could not reliably be indicated by the delta Ct (hsa-miR-23a-3p - hsa-miR-451a) microRNA quantification method. This is an important contrast with serum samples, where hemolysis can be quantified accurately using this microRNA

method (8,20). In our comparison study, the majority of the plasma samples were falsely classified as showing hemolysis by the delta Ct method. This was predominantly due to reduced levels of the housekeeping microRNA hsa-miR-23a-3p, rather than alterations in the levels of the direct hemolysis marker hsa-miR-451a. In our time-course study, delta Ct hemolysis levels in the plasma samples increased then plateaued during a period of ongoing hemolysis (which was indicated by sample inspection and spectrophotometric measurement of hemoglobin absorbance). Normalization of hsa-miR-23a-3p levels to those of hsa-miR-451a, as required for other housekeeping microRNAs in stored plasma samples, would not be appropriate when using the delta Ct (hsa-miR-23a-3p - hsa-miR-451a) method for hemolysis quantification.

Processing of plasma samples for ctDNA quantification typically involves a double centrifugation process, which includes a high-speed ($\geq 10,000g$) second spin, designed to remove background genomic DNA contamination (32-34). Our experimental findings and re-analysis of published data (42) show that such a two-step process significantly alters the profile of circulating microRNAs detected in plasma samples, compared with plasma samples processed by low-speed centrifugation only and also with serum samples. These findings are consistent with a previous global microRNA profiling study showing that the two-step method, including high-speed centrifugation, resulted in lower microRNA levels in plasma when compared with serum (43). Our analyses indicate that the alterations in plasma microRNA levels produced by high-speed centrifugation are highly variable and unpredictable. The microRNAs affected most may predominantly be present in relatively large structures in the plasma, such as platelets or large extracellular vesicles, that are removed by the high-speed centrifugation step (42).

Taken together, our data allow us to make recommendations for the design of biological studies linked to clinical trials, and/or retrospective studies of samples in biorepositories, where quantification of circulating microRNAs is a requirement. Our findings indicate that serum is the optimum biospecimen for quantifying circulating microRNAs, as it is subject to the least pre-analytical technical variation. The optimal approach for studies measuring both circulating microRNAs and ctDNA would therefore be to collect both serum (for microRNAs) and plasma (for ctDNA). If only plasma can be collected, then immediate processing is recommended, in order to avoid the technical variations that affect microRNA levels after storage and/or transport of plasma at room temperature. We further recommend that such plasma processing should involve a low-speed centrifugation step only, followed by aliquoting the supernatant into parallel samples prior to storage at -80°C. Such aliquots could then be used directly for microRNA quantification, or subjected to a further high-speed ($\geq 10,000g$) centrifugation step to optimise ctDNA retrieval (42).

For retrospective analysis of microRNA levels in plasma samples from biorepositories, particularly those prepared for ctDNA extraction, we recommend normalization of housekeeping microRNA levels using cel-miR-39-3p (correcting for technical differences in RNA recovery) and hsa-miR-451a (correcting for the effects of hemolysis), and also the use of multiple housekeeping microRNAs for referencing the levels of test microRNAs. These steps will improve the likelihood of avoiding biologically inconsistent or even contradictory results, as has recently been highlighted for various urological tumors (7). Despite these steps, our data indicate that the results of retrospective microRNA quantification work in such plasma samples should be interpreted with caution.

In our opinion, these technical considerations are of vital importance to the design of future clinical trials and/or retrospective studies of biobank samples. Our recommendations are practicable and scalable. They will help future-proof clinical studies in which quantification of circulating microRNAs is a component.

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Legends to Figures and Tables

Figure 1. Levels of individual microRNAs in samples from the four types of blood collection tube in the comparison study. The plots show Ct values for: A) the exogenous non-human spike-in microRNA cel-miR-39-3p; B)-C) the microRNAs hsa-miR-23a-3p and hsa-miR-451a used for hemolysis assessment; and D)-F) the three established endogenous housekeeping microRNAs hsa-miR-30b-5p, hsa-miR-30c-5p, hsa-miR-191-5p. Key to blood collection tubes: GSS = gel separator serum (green boxes); EP = EDTA plasma (blue); SDP = Streck DNA plasma (red); SRP = Streck RNA plasma (grey). Bar = median, box = interquartile range; whiskers = full range of data.

Figure 2. Levels of hemolysis in plasma samples from the time-course study. A) Appearance of the plasma samples derived from the SDP and EP tubes taken from two healthy control subjects (CONT_001 and CONT_002) at the six time-points [day (d) 0, d2, d4, d7, d10 and d14]. The darker the sample color, the greater the degree of hemolysis. B) Spectrophotometric analysis of the plasma samples in A), showing mean absorbance of free hemoglobin at 414nm (A_{414}). The lower dotted line indicates the threshold (0.2) above which samples were classified as hemolyzed, while the upper dotted line (3.0) indicates the upper limit of detection of the spectrophotometer. C) Hemolysis assessment by delta Ct (hsa-miR-23a-3p – hsa-miR-451a) values. The dotted line indicates the Ct value threshold (8) above which samples were classified as hemolyzed. Key: EP = EDTA plasma (blue lines); SDP = Streck DNA plasma (red lines). Error bars = SD.

Figure 3. MicroRNA levels in plasma samples from the time-course study. Graphs show Ct values for the plasma samples derived from the SDP and EP tubes taken from two healthy control subjects (CONT_001 and CONT_002) at the six time-points [day (d) 0, d2, d4, d7, d10 and d14]. Panels A)-F) show raw Ct values (solid lines) for: A) the exogenous non-human spike-in microRNA cel-miR-39-3p; B)-C) the microRNAs hsa-miR-23a-3p and hsa-miR-451a used for hemolysis assessment; and D-F) the three established endogenous housekeeping microRNAs hsa-miR-30b-5p, hsa-miR-30c-5p and hsa-miR-191-5p. Panels G)-I) show the hemolysis-corrected Ct values (dotted lines) for the three endogenous housekeeping microRNAs shown in D)-F). Key: EP = EDTA plasma (blue lines); SDP = Streck DNA plasma (red lines). Note that error bars are not visible as they fall within each respective data-point.

Figure 4. Comparison of methods of hemolysis quantification. A)-B) Measurement of hemolysis in samples from the comparison study using: mean absorbance of free hemoglobin at 414nm (A_{414}), as assessed by spectrophotometry (A); and delta Ct (hsa-miR-23a-3p - hsa-miR-451a) values (B). C)-D) Raw Ct hsa-miR-451a values (C) and delta Ct (hsa-miR-23a-3p - hsa-miR-451a) values (D) in 32 EP and SDP samples from both the comparison and time-course studies, identified by spectrophotometry as non-hemolyzed (yellow box; A_{414} values ≤ 0.2) or hemolyzed (orange box; A_{414} values > 0.2). E)-F) Associations of raw Ct hsa-miR-451a values (E) and delta Ct (hsa-miR-23a-3p - hsa-miR-451a) values (F) with A_{414} absorbance for the 32 plasma samples analyzed in C) and D), including the lines of best fit. Key: GSS = gel separator serum (green box); EP = EDTA plasma (blue); SDP = Streck DNA plasma (red). Bar = median, box = interquartile range; whiskers = full range of data. Hemolysis thresholds are indicated by the dotted lines, as defined in the legend to Figure 2.

Figure 5. Effect of centrifugation speed on microRNA levels in plasma samples. A) Ct values (cel-miR-39-3p-normalized) for five endogenous microRNAs (hsa-miR-30b-5p, hsa-miR-30c-5p, hsa-miR-191-5p, hsa-miR-23a-3p and hsa-miR-451a) in the GSS, EP #1 and EP #2 samples from patient CUB_008; B) Individual Ct values (cel-miR-39-3p-normalized) for the five endogenous microRNAs assessed in A). Key for A)-B): GSS = gel separator serum (green box); EP #1 = EDTA plasma (low-speed centrifugation only, light blue); EP #2 = EDTA plasma (including high-speed centrifugation, blue). C) Mean Ct values for 106 endogenous microRNAs in EP samples processed using different centrifugation speeds (42). D)-E) Individual raw Ct values for four of the five endogenous microRNAs from panels A) and B) (D) and all 106 microRNAs (E) in EP samples processed using different centrifugation speeds. F) Differences in Ct values for all 106 microRNAs in EP samples centrifuged at 10,000g vs. 2,000g. Key for C)-F): EP 1,000g = EDTA plasma samples with low-speed centrifugation at 1,000 g (light-blue); EP 2,000g = EDTA plasma samples with low-speed centrifugation at 2,000 g (mid-blue); EP 10,000g = EDTA plasma samples with high-speed centrifugation at 10,000 g (dark-blue). In panels A) and C), bar = median, box = interquartile range; whiskers = full range of data.

Table 1. MicroRNA levels in each of the four blood collection tubes from the comparison study. Comparison of Ct values and SD values between the different sample types. Ct values for sample sets were defined as being *suboptimal* if mean microRNA expression values were ≥ 2 Cts greater than the set with the lowest Ct value. SD values for sample sets were defined as *suboptimal* if they were ≥ 2 . For both Ct and SD assessment, samples that were not suboptimal were deemed to be *satisfactory*. Key to blood collection

tubes: GSS = gel separator serum; EP = EDTA plasma; SDP = Streck DNA plasma; SRP = Streck RNA plasma.

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Figure 1

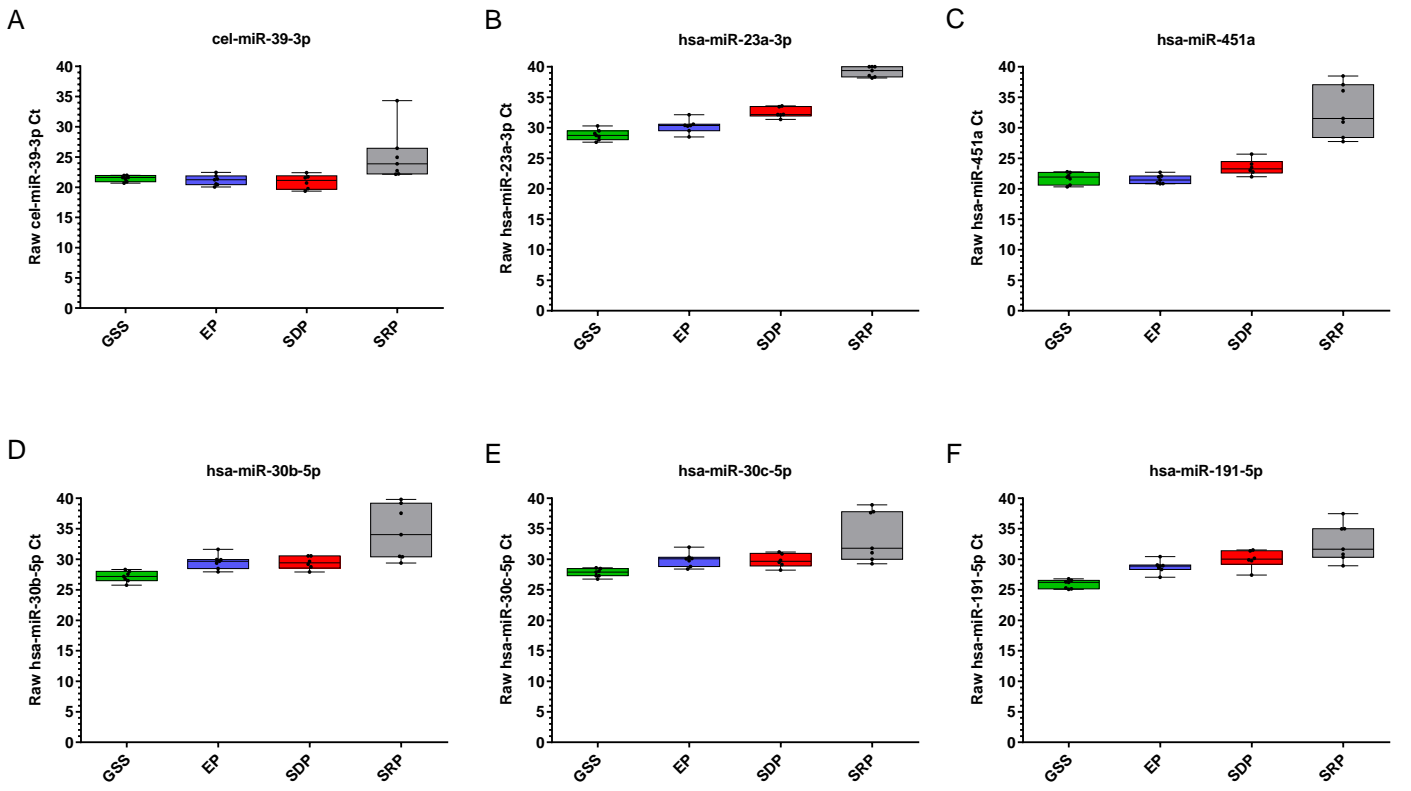
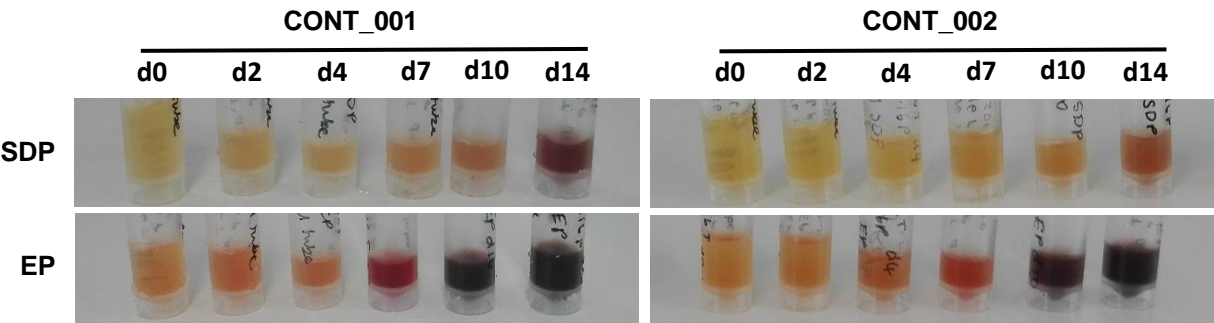
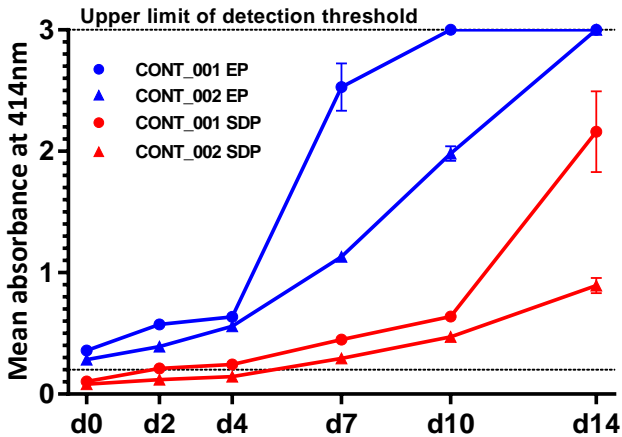


Figure 2

A



B



C

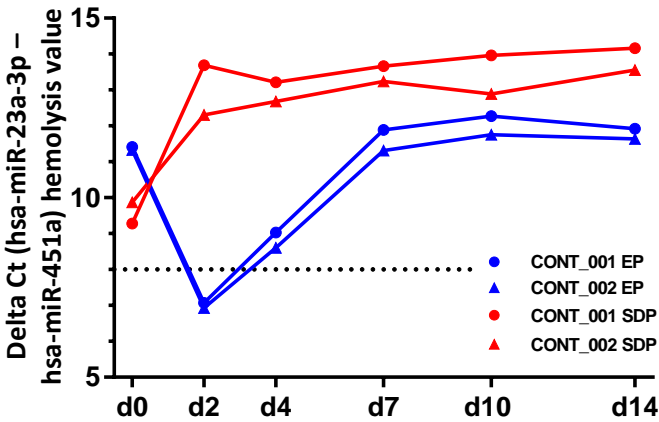


Figure 3

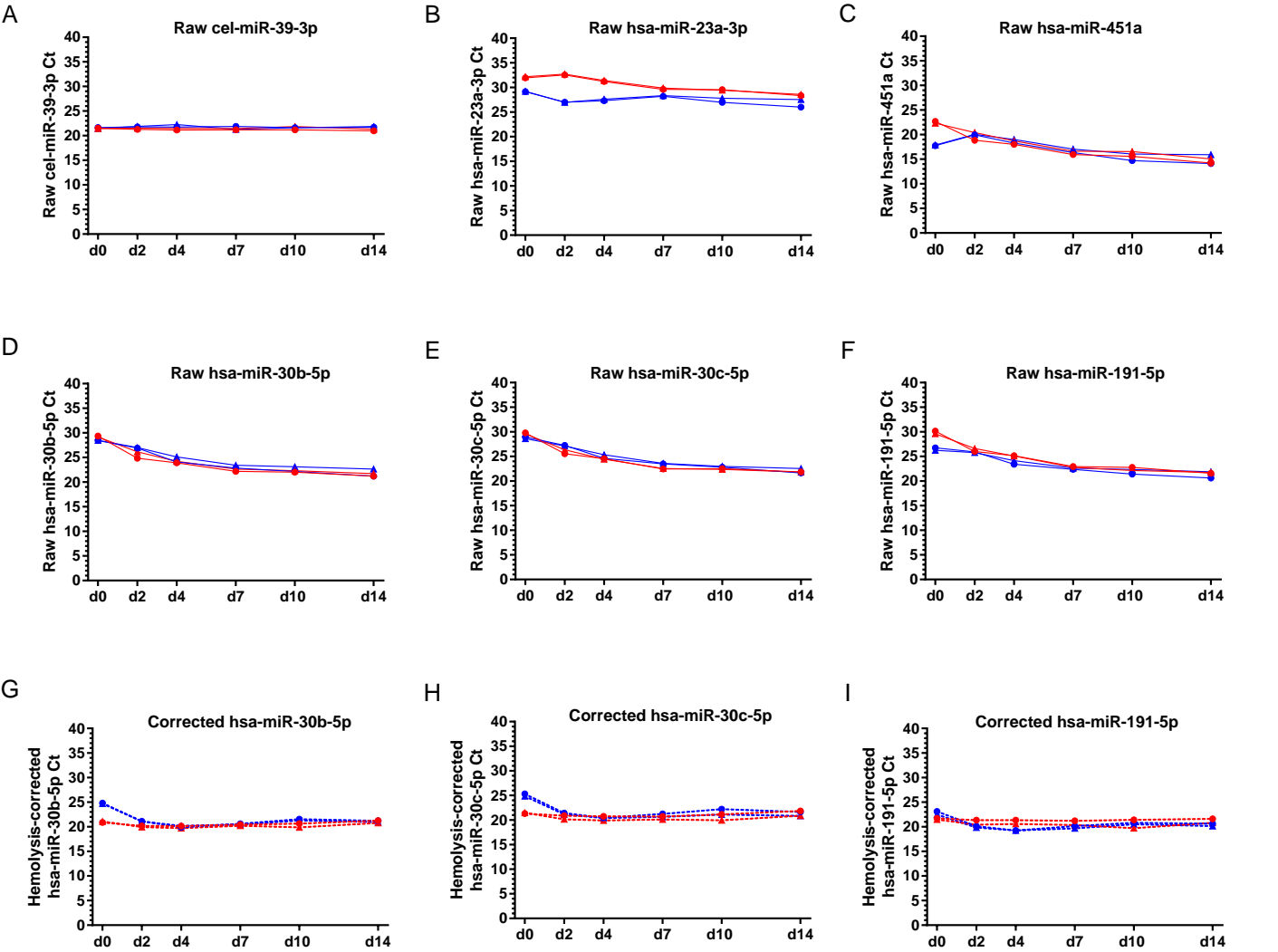
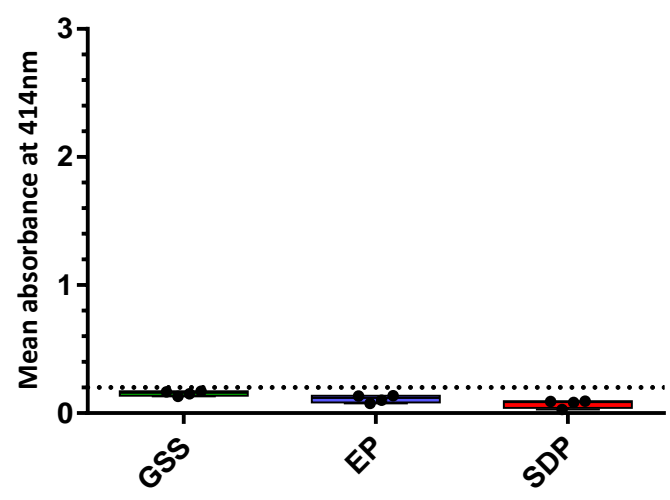
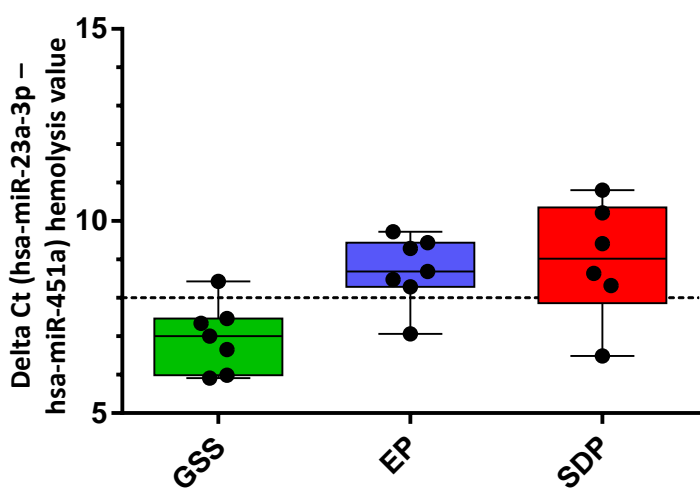


Figure 4

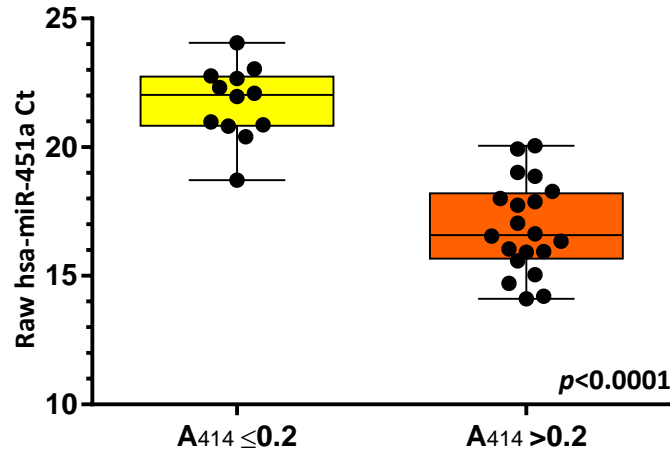
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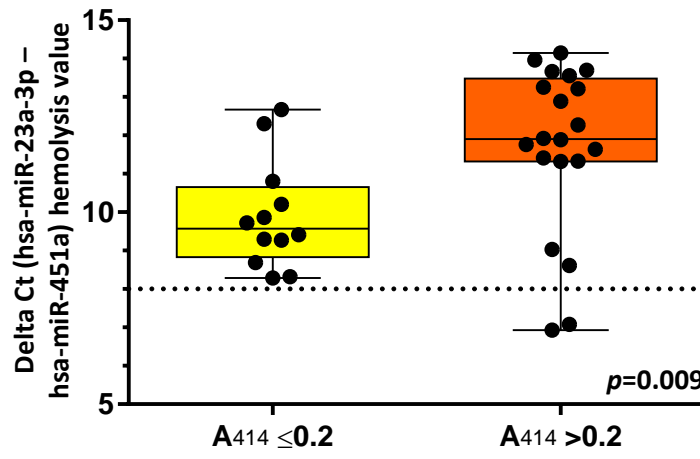
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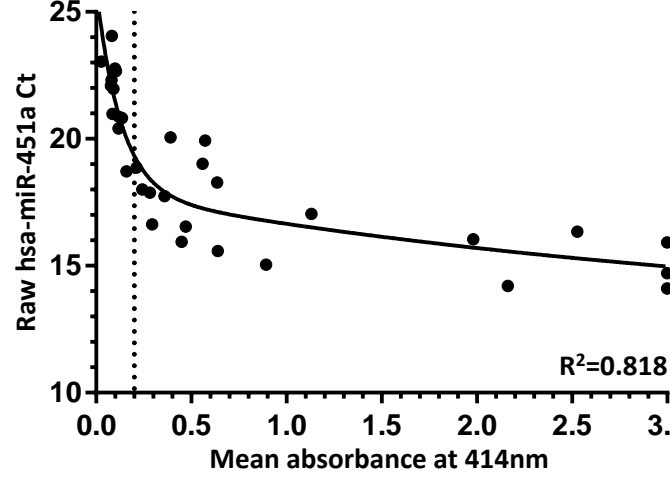
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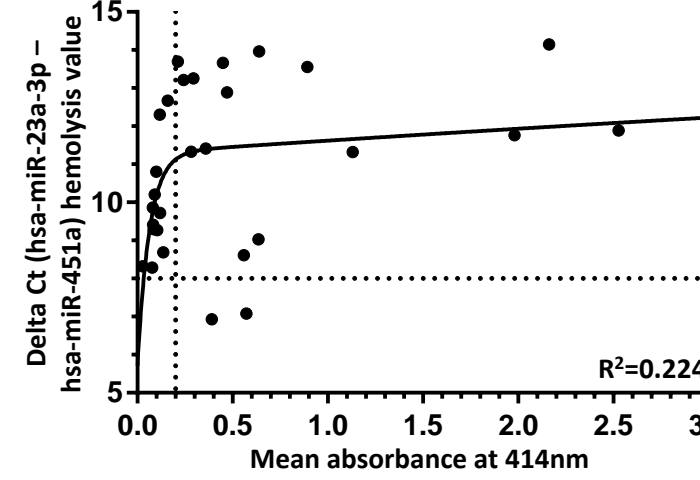
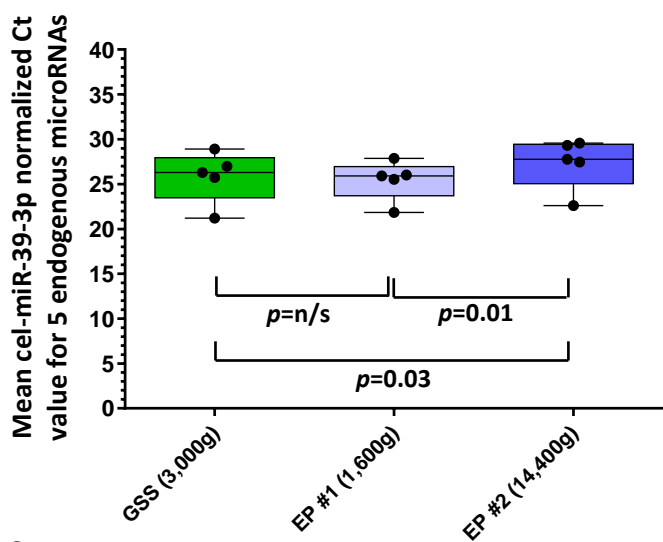
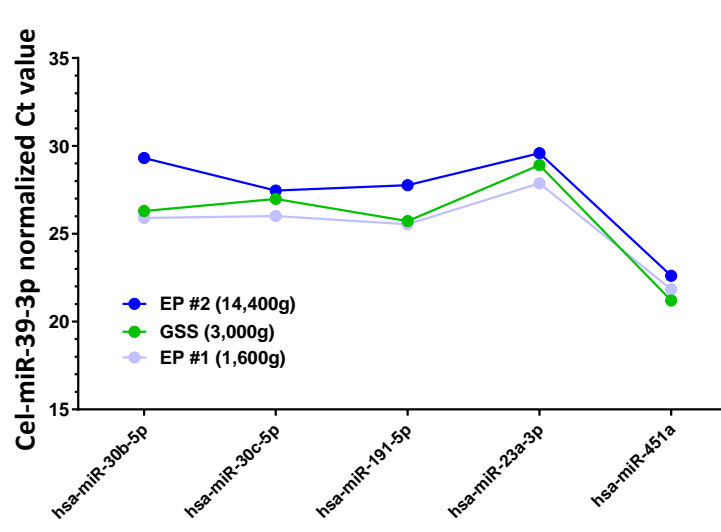


Figure 5

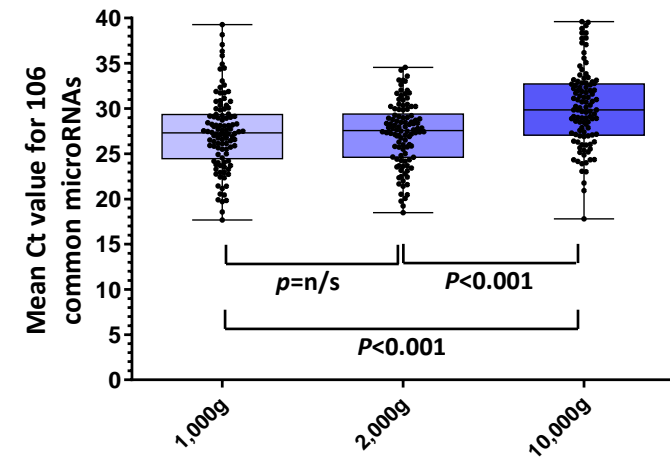
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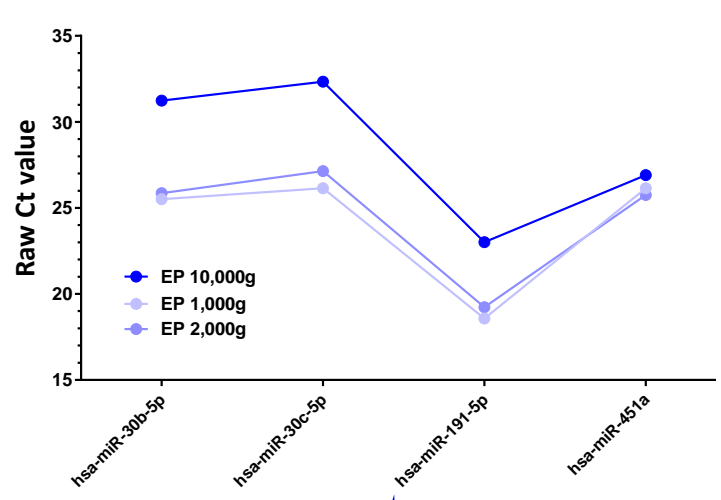
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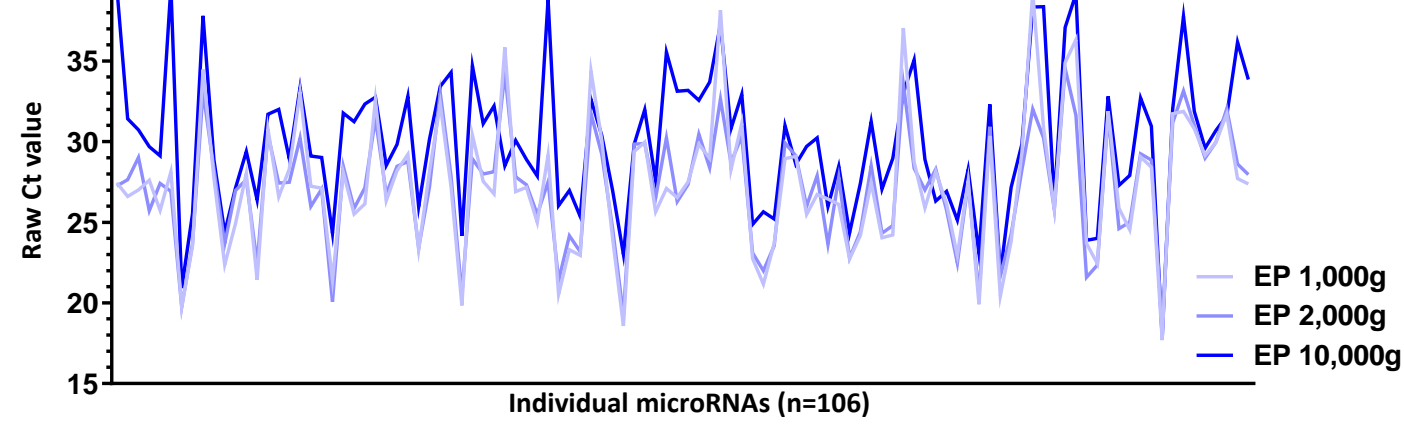
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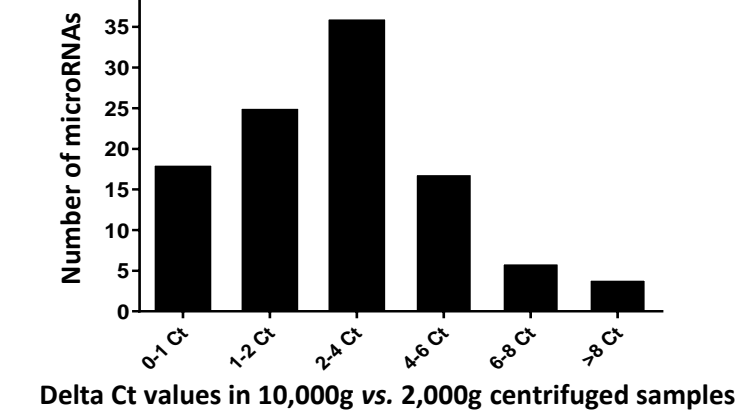


Table 1

microRNA	Role	GSS			EP			SDP			SRP		
		Mean Ct value	SD	Conclusion	Mean Ct value	SD	Conclusion	Mean Ct value	SD	Conclusion	Mean Ct value	SD	Conclusion
cel-miR-39-3p	Exogenous normalization	21.43	0.49	Satisfactory	21.13	0.85	Satisfactory	20.91	1.19	Satisfactory	25.23	4.29	Suboptimal
hsa-miR-30b-5p	Endogenous normalization	27.16	0.90	Satisfactory	29.57	1.18	Suboptimal	29.44	1.04	Suboptimal	34.41	4.45	Suboptimal
hsa-miR-30c-5p	Endogenous normalization	27.77	0.68	Satisfactory	29.93	1.16	Suboptimal	29.73	1.11	Suboptimal	33.78	4.16	Suboptimal
hsa-miR-191-5p	Endogenous normalization	25.91	0.72	Satisfactory	28.76	1.00	Suboptimal	30.02	1.49	Suboptimal	32.74	3.10	Suboptimal
hsa-miR-23a-3p	Endogenous normalization (hemolysis)	28.84	0.89	Satisfactory	30.26	1.10	Satisfactory	32.47	0.86	Suboptimal	39.20	0.84	Suboptimal
hsa-miR-451a	Hemolysis assessment	21.73	0.96	Satisfactory	21.55	0.72	Satisfactory	23.50	1.27	Satisfactory	32.90	4.30	Suboptimal